





#4 D3 6-14

I HEREBY CERTIFY that annexed hereto is a true copy of documents filed in connection with the following patent application:

Application No.

PCT/IE99/00036

Date of Filing

7 May 1999

Applicant

SALVIAC LIMITED, an Irish company of 39-40

Upper Mount Street, Dublin 2, Ireland.

Dated this  $i \rightarrow$  day of August 2001.



Coherly

An officer authorised by the Controller of Patents, Designs and Trademarks.

## HOME COPY

See Notes to the request form

# **PCT**

## **REQUEST**

The undersigned requests that the present international application be processed according to the Patent Cooperation Treaty.

For receiving Office use only

PCT/E 9 9 / 0 0 0 3 6
International Application No.

- 7 MAY 1999
International Filing Date 10 7. 05. 99)

PRISH PATENICS OFFICE

POT INTERNATIONAL APPLICATION

Name of receiving Office and PCT International Application

	Applicant's or agent's file reference if desired) (12 characters maximum) SALVO9/C/WO						
Box No. I TITLE OF INVENTION							
"Improved Filter Element for Embolic Protection Device"							
Box No. II APPLICANT							
Name and address: (Family name followed by given name: for a le designation. The address must include postal code and name of count address indicated in this Box is the applicant's State (that is, country) of residence is indicated below.)	gal entity, full official y. The country of the of residence if no State  This person is also inventor.						
SALVIAC LIMITED	Telephone No.						
39-40 Upper Mount Street							
Dublin 2	Facsimile No.						
Ireland							
	Teleprinter No.						
State (that is, country) of nationality:	State (that is, country) of residence:						
IE	IE .						
This person is applicant all designated for the purposes of:	States except the United States the States indicated in the Supplemental Box						
Box No. III · FURTHER APPLICANT(S) AND/OR (FURTH)	ER) INVENTOR(S)						
Name and address: (Family name followed by given name: for a le designation. The address must include postal code and name of country address indicated in this Box is the applicant's State (that is country) to residence is indicated below.)  VALE, David 26 The Stiles Road Clontarf Dublin 3 Ireland	This person is:  This person is:  applicant only  X applicant and inventor  inventor only (If this check-box is marked, do not fill in below.)						
State (that is, country) of nationality:	State (that is. country) of residence:						
IE	IE						
This person is applicant all designated all designated to the purposes of:	States except the United States the States indicated in the Supplemental Box						
X Further applicants and/or (further) inventors are indicated on	a continuation sheet.						
Box No. IV AGENT OR COMMON REPRESENTATIVE:	OR ADDRESS FOR CORRESPONDENCE						
The person identified below is hereby/has been appointed to act on of the applicant(s) before the competent International Authorities as	behalf X agent Common representative						
Name and address: (Family name followed by given name: for a legal entity, full official designation. The address must include postal code and name of country.)  + 353 1 2883877							
O'BRIEN, John A: WELDON, Michael J							
c/o John 4 O'Brien & Associates,	Facsimile No. + 353 1 2993878						
Third Floor, Duncairn House,	many and a second secon						
<pre>14 Carysfort Avenue. Blackrock.</pre>	Teleprinter No.						
County Dubin,							
Treland.	The state of the s						
Address for correspondence: Mark this check-box where no space above is used instead to indicate a special address to wh	agent or common representative is has been appointed and theinches correspondence should be sent.						

Sheer	Sin	. 2 .	
111001	. v().	• • • • •	

Continuation of Box No. III FURTHER APPLICANT(S) AND/OR (FURTHER) INVENTOR(S)						
	his sheet should not be included in the request.					
Name and address: (Family name followed by given name: for a designation. The address must include postal code and name of cour address indicated in this Box is the applicant's State (that is, country) of residenc: is indicated below.)  BRADY, Eamon 12 Karol avenue Elphin County Roscommon Ireland	legal entity, full official lity. The country of the of residence if no State  This person is:  applicant only  X applicant and inventor  inventor only (If this check-box is marked, do not fill in below.)					
State (that is, country) of nationality:	State (that is, country) of residence:					
This person is applicant all designated all designated	States except the United States the States indicated in tes of America only the Supplemental Box					
Name and address: (Family name followed by given name: for a ladesignation. The address must include postal code and name of count address indicated in this Box is the applicant's State (that is, country) of residence is indicated below.)						
State (that is, country) of nationality:	State (that is, country) of residence:					
This person is applicant all designated all designated for the purposes of:  Name and address: (Family name followed by given name: for a led designation. The address must include postal code and name of count address indicated in this Box is the applicant's State (that is, country) of residence is indicated below.)	es of America only the Supplemental Box					
of residence is indicated below.)	This person is:  applicant only  applicant and inventor  inventor only (If this check-box is marked, do not fill in below.)					
State (that is, country) of nationality:	State (that is, country) of residence:					
This person is applicant all designated all designated for the purposes of:	States except the United States the States indicated in the Supplemental Box					
Name and address: (Family name followed by given name: for a le designation. The address must include postal code and name of count address indicated in this Box is the applicant's State (that is country) of of residence is indicated below.)	gal entity, full official y. The country of the off residence if no State  This person is:  applicant only  applicant and inventor  inventor only if this check-box is marked, so not fill in below.)					
State (that is, country) of nationality:	State (that is country) of residence: (6)					
This person is applicant all designated the United States all designated the United States	States except the United States the States indicated in the Supplemental Box					
Further applicants and/or (further) inventors are indicated on	another continuation sheet, and which we have the same of the same					

Box	10. V	DESIGNATION OF STATES							
The to	How	ing designations are hereby made under Rule 4.9(a	) incl	rk the	applicable check-hoxes, at least one must be marked)				
Regio		3 3	•						
N.		•	ı. I.S	Leson	no. MW Malawi, SD Sudan, SZ Swazifand, UG Uganda				
0.3		ZW Zimbabwe, and any other State which is a Conti	ractin	ig Stat	e of the Harare Protocol and of the PCT				
· <b>3</b>	EΑ	Moldova, RU Russian Federation, TJ Tajikistan, T.	BY M Tu	Belan irkmei	is. KG Kyrgyzstan, KZ Kazakhstan, MD Republic of histan, and any other State which is a Contracting State				
[2]	c n	of the Eurasian Patent Convention and of the PCT		. r.c	installand and Liashtanetain CV Cyngus DE Com-				
K	EP	EP European Patent: AT Austria, BE Belgium, CH and LI Switzerland and Liechtenstein, CY Cyprus, DE Germany, DK Denmark, ES Spain, FI Finland, FR France, GB United Kingdom, GR Greece, IE Ireland, IT Italy, LU Luxembourg, MC Monaco, NL Netherlands, PT Portugal, SE Sweden, and any other State which is a Contracting State of the European Patent Convention and of the PCT							
OA OAPI Patent: BF Burkina Faso, BJ Benin, CF Central African Republic, CG Congo, CI Côte d'Ivoire, CM Cameroon, GA Gabon, GN Guinea, GW Guinea-Bissau, ML Mali, MR Mauritania, NE Niger, SN Senegal, TD Chad. TG Togo, and any other State which is a member State of OAPI and a Contracting State of the PCT (if other kind of protection or treatment desired, specify on dotted line)									
		tent (if other kind of protection or treatment desired, specify		ottea ti	ne):				
<b>⊠</b>		Albania	X		Lesotho				
⊠		Armenia	$\boxtimes$		Lithuania				
$\boxtimes$		Austria	$\boxtimes$	LU	Luxembourg				
	ΑU	Australia	$\boxtimes$	_	Latvia				
$\boxtimes$	ΑZ	Azerbaijan	$\square$	MD	Republic of Moldova				
$\boxtimes$	BA	Bosnia and Herzegovina	$\boxtimes$	MG	Madagascar				
$\boxtimes$	BB	Bashados	$\boxtimes$	MK	The former Yugoslav Republic of Macedonia				
$\boxtimes$	ВG	Bulgaria							
<b>⊠</b>	BR	Brazil	$\boxtimes$	MN	Mongolia				
$\boxtimes$	BY	Belarus	$\boxtimes$	мw	Malawi				
<b>⊠</b>		Canada	$\overline{\mathbb{N}}$		Mexico				
⊠ ⊠		and LI Switzerland and Liechtenstein	$\boxtimes$		Norway				
		China	$\boxtimes$		New Zealand				
		Cuba	$\boxtimes$		Poland				
		Czech Republic	$\boxtimes$		Portugal				
× ×		Germany . and utility model	$\boxtimes$		Romania				
		-			Russian Federation				
$\overline{\Delta}$		Denmark and .utility model	$\boxtimes$						
X		Estonia	$\boxtimes$	SD	Sudan				
X	ES	Spain		SE	Sweden				
$\boxtimes$	FI	Finland	$\boxtimes$	SG	Singapore				
$\boxtimes$	GB	United Kingdom	$\boxtimes$	SI	Slovenia				
N N	_	Grenada	$\boxtimes$		Slovakia				
	GE	Georgia	Δ.	SL	Sierra Leone				
	GH	Ghana	$\boxtimes$	TJ	Tajikistan				
$\boxtimes$		Gambia	$\boxtimes$	TM	Turkmenistan				
	HR	Croatia	$\boxtimes$	TR	Turkey				
	HU	Hungary	$\square$	TT	Trinidad and Tobago				
	ID	Indonesia	$\boxtimes$	UА	Ukraine				
$\Sigma$	IL	Israel	$\boxtimes$	UG	Uganda				
□ □	IN	India	$\boxtimes$	US	United States of America				
Image: second color in the secon	IS	Iceland							
	JP	Japan	$\boxtimes$	UΖ	Uzbekistan				
図	KE	Kenya	$\square$	VN	Viet Nam				
×	KG	Kyrgyzstan	$\boxtimes$		Yugoslavia				
	KP	Democratic People's Republic of Korea	$\overline{\boxtimes}$		Zimbabwe				
	rca -	Democratic respite 5 Republic of Rolles	_						
(5)	L/D	Republic of Korea	Che a no	cx-co: tional	ces reserved for designating States (for the purposes of patent) which have become party to the PCT after				
		Kazakhstan		ance o	i this sheet:				
			$\square$	74	South Africa				
	LC	Saint Lucia							
		Sri Lanka		AC	United Arab Emerates				
N N		Liberia							
Preca	ution	ary Designation Statement: In addition to the designa	tions	made.	above, the applicant also makes under Rule 4.9(b) all other				

Precautionary Designation Statement: In addition to the designations made above, the applicant also makes under Rule 4.9(b) all other designations which would be permitted under the PCT except any designation(s) indicated in the Supplemental Box as being excluded from the scope of this statement. The applicant declares that those additional designations are subject to confirmation and that any designation which is not confirmed before the expiration of 15 months from the priority date is to be regarded as withdrawn by the applicant at the expiration of that time limit. (Confirmation of a designation consists of the filling of a notice specifying that designation and the payment of the designation and confirmation fees. Confirmation must reach the receiving Office within the 15-month (time limit).

Sheet	\u0	4
11.000		

Fling date of earlier application (day,month/year)  item (1)  The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application is the receiving Office)  The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application was filed with the Office which for the purposes of the present international application is the receiving Office)  * Where the earlier application is an ARPO application is the readjugation in the Supplemental Bo  Further priority claims are indicated in the Supplemental Bo  Where earlier application is:  Tregional Office international application:  regional Office  receiving Office  international Bureau a certified copy of the earlier application is the receiving Office) identified above as item(s):  * Where the earlier application is an ARPO application is the receiving Office in the Supplemental Bo							
item (1)  Item (2)  Item (3)  The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application was filed with the Office which for the purposes of the present international application is the receiving Office) identified above as item(s):							
item (2)  item (3)  The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application was filed with the Office which for the purposes of the present international application is the receiving Office) identified above as item(s):							
The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application was filed with the Office which for the purposes of the present international application is the receiving Office) identified above as item(s):							
The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application was filed with the Office which for the purposes of the present international application is the receiving Office) identified above as item(s):							
The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application was filed with the Office which for the purposes of the present international application is the receiving Office) identified above as item(s):							
The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application was filed with the Office which for the purposes of the present international application is the receiving Office) identified above as item(s):							
The receiving Office is requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) (only if the earlier application was filed with the Office which for the purposes of the present international application is the receiving Office) identified above as item(s):							
purposes of the present international application is the receiving Office) identified above as item(s):							
Where the earlier application is an ARIPO application is is mandaton; to indicate in the Sunday of t							
• Where the earlier application is an ARIPO application, it is mandatory to indicate in the Supplemental Box at least one country party to the Particonvention for the Protection of Industrial Property for which that earlier application was filed (Rule 4.10(b)(ii)). See Supplemental Box.							
Box No. VII INTERNATIONAL SEARCHING AUTHORITY							
Choice of International Searching Authority (ISA) (if two or nore International Searching Authorities are competent to carry out the international search, indicate the Authority chosen: the two-letter code may be used):  Request to use results of earlier search; reference to that search (if an earlier search has been carried out by or requested from the International Searching Authority)  Date (day/month/year)  Number  Country (or regional Office							
to regional Once							
ISA /EP  Box No. VIII CHECK LIST: LANGUAGE OF FILING							
the following number of sheets:							
request : 4 1. X fee calculation sheet							
description (excluding sequence listing part):  11   2.							
sequence listing part) : 11   3.							
abstract : 1 5. priority document(s) identified in Box No. VI as item(s):							
drawings : 9 6. ☐ translation of international application into (language):							
sequence listing part 7   separate indications concerning deposited microorganism or other biological metastical							
of description : Separate indeations echicering deposited interoofganism of other biological material and/or amino acid sequence listing in computer readable form							
Total number of sheets: 27 9. □ other (specify):							
Figure of the drawings which should accompany the abstract: Fig. 18  Language of filing of the international application: English							
Box No. IX SIGNATURE OF APPLICANT OR AGENT							
Next to each signature, indicate the name of the person signing and the capacity in which the person signs (if such capacity is not obvious from reading the request).							
Loluthia:							
John O'Brien							
For receiving Office use only							
1. Date of actual receipt of the purported = 7 MAY 1999 2. Drawings: 107.05.99							
3. Corrected date of actual receipt due to later but timely received papers or drawings completing the purported international application:							
4. Date of timely receipt of the required corrections under PCT Article 11(2):							
5. International Searching Authority ISA / EP 6. Transmittal of search copy delayed until search fee is paid.							
For International Bureau use only							
Dage of receipt of the record copy by the International Bureau:							

#### "IMPROVED FILTER ELEMENT FOR EMBOLIC PROTECTION DEVICE"

This invention relates to a filter element for a transcatheter embolic protection device.

5

10

15

20

#### **Introduction**

The invention is particularly concerned with filter elements for transcatheter embolic protection devices of the type described in our previously filed PCT Patent Application No. PCT/IE98/00093 the contents of which are incorporated herein by reference. One type of such embolic filter essentially comprises a filter body mounted on an associated collapsible support frame which can be collapsed against the guide wire by means of a catheter for deployment of the filter through a patient's vascular system. Upon retraction of the catheter the support frame and filter body expand outwardly from the guidewire across a blood vessel within which the filter is positioned to filter blood flowing through the blood vessel.

A practical problem that arises with filter elements of such embolic protection devices is that they should be able to accommodate blood vessels of different diameter as it would be impractical to manufacture a large range of filters each of different size to accommodate all possible diameters of blood vessel. To provide flexibility and accommodate a range of vessel sizes with a given size of filter a relatively soft and elastic filter body material can be used. It is, however, important that the filter when deployed maintains its shape during use and to prevent distortion or collapsing of the filter body in use. Because of this and also the need for adequate strength in the body material, the walls of the filter body tend to be relatively thick. This presents a problem in that the filter then has a relatively large crossing profile when in the collapsed deployment position, which is undesirable.

30

25

The present invention is directed towards overcoming these problems.

## Statements of Invention

5

10

15

20

30

According to the invention there is provided a collapsible filter element comprising:

a collapsible filter body which is movable between a collapsed stored position for movement through a vascular system and an expanded position for extension across a blood vessel such that blood passing through the blood vessel is delivered through the filter element;

an inlet end of the filter body having one or more inlet openings sized to allow blood and embolic material enter the filter body;

an outlet end of the filter body having a plurality of outlet openings sized to allow through-passage of blood, but to retain embolic material within the filter body;

wherein the filter body has a first relatively stiff portion and a second relatively soft portion for engaging a circumferential vessel wall.

Ideally the first portion has a larger wall thickness than the wall thickness of the second portion.

In a particularly preferred embodiment of the invention the filter body is of composite construction.

In one embodiment of the invention, the filter body comprises a proximal body section and a distal body section, one of which forms said stiff first portion and the other forming the soft second portion.

Preferably the proximal body section forms the soft second portion.

In a further embodiment, the filter body comprises a proximal body section and a distal body section interconnected by an intermediate body section, one or both of the proximal body section and the intermediate body section forming the soft second portion, the distal body section forming the stiff first portion.

In another embodiment the proximal body section has a ribbed outer surface.

Preferably a plurality of spaced-apart longitudinal ribs are provided on the proximal section.

In another embodiment the proximal body section includes corrugations.

## Brief Description of the Drawings

The invention will be more clearly understood by the following description of some of the embodiments thereof, given by way of example only, with reference to the accompanying drawings, in which: -

20

15

5

Fig. 1 is partially sectioned elevational view an embolic protection device;

Fig. 2 is a schematic sectional elevational view of the embolic protection device of Fig. 1;

25

30

Fig. 3 is a detail sectional view of portion of the device of Fig. 1;

Fig. 4 is a longitudinal cross sectional view of the device of Fig. 1;

Fig. 5 is a cross sectional view of a distal end of the device of Fig. 1;

		_		_								
T:-	_		_	view		41	1:	A A	•	T- '	~	
r 1O	n	16	2	VIEW	an	TNP	פתוו	$\Delta - \Delta$	าก	HIG	<b>→</b> :	,
4 15.	v	IJ	u	4 TC 44		$u_{1}$	1111	$\Delta$	111	1 12.	J.	

Fig. 7 is a perspective view of a filter body of the device of Figs. 1 to 6;

Fig. 8 is a side elevational view of the filter body of Fig. 7;

Fig. 9 is a view on a proximal end of the filter body;

Fig. 10 is a perspective view of a support frame of the device of Figs. 1 to 6;

Fig. 11 is a side elevational view of the support frame;

Fig. 12 is a perspective view illustrating the manufacture of the support frame;

Fig. 13 is a view of the support frame and filter element assembly;

Fig. 14 is a longitudinal cross sectional view of a filter body according to the invention;

Fig. 15 is a longitudinal cross sectional view of another filter body according to the invention;

Fig. 16 is a longitudinal cross sectional view of a further filter body according to the invention;

Fig. 17 is a schematic perspective view of a filter element according to another aspect of the invention; and

Fig. 18 is another schematic perspective view of a filter element of the invention.

10

5

15

20

25

30

#### **Detailed Description**

10

15

20

25

30

Referring to Figs. 1 to 13 there is illustrated an embolic protection device as described in our co-pending Application PCT/IE98/00093 indicated generally by the reference number 100. The device 100 has a guidewire 101 with a proximal end 102 and a distal end 103. A tubular sleeve 104 is slidably mounted on the guidewire 101. A collapsible filter 105 is mounted on the sleeve 104, the filter 105 being movable between a collapsed stored position against the sleeve 104 and an expanded position as shown in the drawings extended outwardly of the sleeve 104 for deployment in a blood vessel.

The sleeve 104 is slidable on the guidewire 101 between a pair of spaced-apart end stops, namely an inner stop 106 and an outer stop which in this case is formed by a spring tip 107 at the distal end 103 of the guidewire 101.

The filter 105 comprises a filter body 110 mounted over a collapsible support frame 111. The filter body 110 is mounted to the sleeve 104 at each end, the body 110 being rigidly attached to a proximal end 112 of the sleeve 104 and the body 110 being attached to a collar 115 which is slidable along a distal end 114 of the sleeve 104. Thus the distal end of the body 110 is longitudinally slidable along the sleeve 104. The support frame 111 is also fixed at the proximal end 112 of the sleeve 104. A distal end 116 of the support frame 111 is not attached to the sleeve 104 and is thus also free to move longitudinally along the sleeve 104 to facilitate collapsing the support frame 111 against the sleeve 104. The support frame 111 is such that it is naturally expanded as shown in the drawings and can be collapsed inwardly against the sleeve 104 for loading in a catheter 118 or the like.

The filter body 105 has large proximal inlet openings 117 and small distal outlet openings 119. The proximal inlet openings 117 allow blood and embolic material

to enter the filter body, however, the distal outlet openings 119 allow through passage of blood but retain undesired embolic material within the filter body.

An olive guide 120 is mounted at a distal end of the sleeve 104 and has a cylindrical central portion 121 with tapered ends 122, 123. The distal end 122 may be an arrowhead configuration for smooth transition between the catheter and olive surfaces. The support frame 111 is shaped to provide a circumferential groove 125 in the filter body 110. If the filter is too large for a vessel, the body may crease and this groove 125 ensures any crease does not propagate along the filter.

5

10

15

20

25

30

Enlarged openings are provided at a proximal end of the filter body 110 to allow ingress of blood and embolic material into an interior of the body 110.

In use, the filter 105 is mounted in a collapsed state within a distal end of the catheter 118 and delivered to a deployment site. When the filter is correctly positioned the catheter 118 is retracted allowing the support frame 111 to expand expanding the filter body 110 across the vessel in which the filter is mounted. Blood and emboli can enter the enlarged openings at a proximal end of the filter body 110. The blood will pass through the net wall, however, the openings or pores in the net are sized so as to retain the embolic material. After use the catheter is delivered along the guidewire 101 and slid over the filter 105 engaging the proximal inlet end 112 first to close the openings and then gradually collapsing the filter body against the sleeve 104 as the catheter 118 advances over the filter 105. Once the filter 105 is fully loaded in the catheter 118, it can then be withdrawn.

It will be noted that a proximal end of the filter is fixed and a distal end of the filter is longitudinally movable along the sleeve to facilitate collapsing of the filter body.

Further, the catheter engages the proximal end of the filter body first thus closing the filter body inlet and preventing escape of embolic material from the filter body as the filter body is being collapsed.

The outer filter body 110 is preferably of a resilient biocompatible elastomeric material. The material may be a polyurethane based material. There are a series of commercially available polyurethane materials that may be suitable. These are typically based on polyether or polycarbonate or silicone macroglycols together with disocyanate and a diol or diamine or alkanolamine or water chain extender.

Examples of these are described in EP-A-461,375 and US 5,621, 065. In addition, polyurethane elastomers manufactured from polycarbonate polyols as described in US 5,254,622 (Szycher) are also suitable.

15

20

25

30

The filter material may also be a biostable polycarbonate urethane article an example of which may be prepared by reaction of an isocyanate, a chain extender and a polycarbonate copolymer polyol of alkyl carbonates. This material is described in our co-pending PCT Application No. IE98/00091, filed November 9, 1998, the entire contents of which are incorporated herein by reference. The filter material may be manufactured from a block and cut into a desired shape. However the filter is preferably formed by dipping a rod of desired geometry into a solution of the material which coats the rod. The rod is then dissolved. The final geometry of the filter may be determined in the dipping step or the final geometry may be achieved in a finishing operation. Typically the finishing operations involve processes such as mechanical machining operations, laser machining or chemical machining.

The filter body is of hollow construction and is formed as described above by dipping a rod in a solution of polymeric material to coat the rod. The rod is then dissolved, leaving a hollow body polymeric material. The rod may be of an acrylic material which is dissolved by a suitable solvent such as acetone.

The polymeric body thus formed is machined to the shape illustrated in Figs. 1 to 13. The final machined filter body comprises an inlet or proximal portion 210 with a proximal neck 212, and outlet or distal portion 213 with a distal neck 214, and an intermediate portion 215 between the proximal and distal portions.

5

The inlet holes 117 are provided in the proximal portion 210 which allow the blood and embolic material to flow into the filter body. In this case the proximal portion 210 is of generally conical shape to maximise the hole size.

10

The intermediate portion 215 is also hollow and in this case is of generally cylindrical construction. This is important in ensuring more than simple point contact with the surrounding blood vessel. The cylindrical structure allows the filter body to come into soft contact with the blood vessel to avoid damaging the vessel wall.

15

The intermediate portion 215 is provided with a radial stiffening means, in this case in the form of a radial strengthening ring or rim 220. The ring 220 provides localised stiffening of the filter body without stiffening the material in contact with the vessel. Such an arrangement provides appropriate structural strength so that line apposition of the filter body to the vessel wall is achieved. It is expected that other geometrics of stiffening means will achieve a similar result.

20

The tubular intermediate portion 215 is also important in maintaining the stability of the filter body in situ to retain captured emboli and to ensure that flow around the filter is minimised. For optimum stability we have found that the ratio of the axial length of the intermediate portion 215 of the filter body to the diameter of the intermediate portion 215 is preferably at least 0.5 and ideally greater than 1.0.

30

25

The collapsible support frame 111 has four foldable arms 290 which are collapsed for deployment and upon release extend outwardly to expand the filter body 110.

The support frame 111 can be manufactured from a range of metallic or polymeric components such as a shape memory alloy like nitinol or a shape memory polymer or a shaped stainless steel or metal with similar properties that will recover from the deformation sufficiently to cause the filter body 110 to open.

5

The support frame may be formed as illustrated in Fig. 12 by machining slots in a tube 291 of shape memory alloy such as nitinol. On machining, the unslotted distal end of the tube forms a distal collar 293 and the unslotted proximal end of the tube forms a proximal collar 294. In use, the distal collar 293 is slidably moveable along the tubular sleeve 104 which in turn is slidably mounted on the guidewire 101 for deployment and retrieval. The proximal collar 294 is fixed relative to the tubular sleeve 104.

10

15

To load the filter the sub assembly of the support frame and filter body is pulled back into the catheter 118 to engage the distal stop 107. The support arms 290 are hinged inwardly and the distal collar 293 moves forward along the tubular sleeve 104. As the support arms 290 enter the catheter 118 the filter body 110 stretches as the filter body collar 115 slides along the tubular sleeve 104 proximal to the olive 120. On deployment, the catheter 118 is retracted proximally along the guidewire 101 initially bringing the collapsed filter assembly with it until it engages the proximal stop 106. The catheter sleeve then begins to pull off the filter freeing the support arms 290 to expand and the filter body apposes the vessel wall.

20

25

30

For retrieval, a retrieval catheter is introduced by sliding it over the guidewire 101 until it is positioned at the proximal end of the filter body and support frame. Pulling the guidewire 101 will initially engage the distal stop 107 with the filter element and begin to pull it into the retrieval catheter. The initial travel into the delivery catheter acts to close the proximal openings of the filter element, thus entrapping the embolic load. As the filter continues to be pulled back the filter

body and the support frame are enveloped in the retrieval catheter. The collapsed filter may then be removed from the patient.

Referring to Fig. 14 there is illustrated a filter element comprising a filter 2 according to the invention. In this case, the filter body 2 has a proximal section 3 and a distal section 4 interconnected by an intermediate section 5. Both the proximal section 3 and the distal section 4 are made from a relatively stiff grade of polyurethane material which enables a low wall thickness to be achieved, thus advantageously minimising the bulk of the filter when it is in a collapsed position so that it has a low crossing profile while at the same time providing adequate strength. The intermediate section 5 is made from a soft elastic grade of polyurethane having good shape memory characteristics which will help the filter maintain the desired expanded shape during use of the filter. This soft portion also allows one filter size to accommodate a range of vessel sizes conforming closely to the vessel wall to prevent blood and embolic material bypassing the filter.

10

15

20

25

30

In the filter body 14 illustrated in Fig. 14 the body is of generally uniform thickness in cross section. However, to achieve any desired variation in the properties of the filter body the thickness may be variable such as in the filter body 10 illustrated in Fig. 15.

Referring to Fig. 16, any required structural properties may also be provided by manufacturing the filter body 20, at least partially from a laminate construction of the same or different materials. In the illustration of Fig. 16 the distal portion 4 and part of the intermediate portion 5 are of twin layer 21, 22 construction. The layers 21, 22 may be of the same of different materials. Keying means, either mechanical or chemical may be provided between the layers. There may be multiple layers and a different layer structure may be provided at any desired locations of the filter body to achieve required properties.

Referring now to Fig. 17 there is shown another filter element according to the invention, indicated generally by the reference 40. The filter element 40 has a filter body 42 of generally similar construction to the filter element described previously with reference to Figs. 1 to 16, the body having a proximal section 43 and a distal section 44 interconnected by an intermediate section 45. In this case, the distal section 44 is of a relatively hard polyurethane material whilst the proximal section 43 and intermediate section 45 are of a softer grade polyurethane material. A number of longitudinal ribs 46 are provided around a circumference of the proximal section 43. Advantageously, this construction facilitates close engagement of an outer circumference of the proximal section 43 against a vessel wall to minimise the risk of embolic material bypassing the filter element 40. An internal spring frame, as described above, urges the proximal section 43 outwardly so that it expands against and closely conforms with the wall of the blood vessel in which the filter element 40 is mounted in use.

15

10

5

Conveniently, the corrugations or ribs 46 allow the proximal section 43 of the filter element 40 to accommodate a wider range of vessel sizes whilst maintaining good contact between the outer circumference of the proximal section 43 and the vessel wall and providing improved web strength and filter body integrity.

20

Referring to Fig. 18 there is illustrated another filter element 50 according to the invention. In this case corrugations 51 are provided for improved apposition and web strength.

25

The invention is not limited to the embodiments hereinbefore described which may be varied in both construction and detail.

#### <u>Claims</u>

A collapsible filter element for a transcatheter embolic protection device, the filter element comprising:

**5** .

a collapsible filter body which is movable between a collapsed stored position for movement through a vascular system and an expanded position for extension across a blood vessel such that blood passing through the blood vessel is delivered through the filter element;

10

an inlet end of the filter body having one or more inlet openings sized to allow blood and embolic material enter the filter body;

15

an outlet end of the filter body having a plurality of outlet openings sized to allow through-passage of blood, but to retain embolic material within the filter body;

20

25

wherein the filter body has a first relatively stiff portion and a second relatively soft portion for engaging a circumferential vessel wall.

- 2. A filter element as claimed in claim 1 wherein the first portion has a larger wall thickness than the wall thickness of the second portion.
- 3. A filter element as claimed in claim 1 or 2 wherein the filter body is of composite construction.
- 4. A filter element as claimed in any preceding claim wherein the filter body comprises a proximal body section and a distal body section, one of which forms said stiff first portion and the other forming the soft second portion.

- 5. A filter as claimed in claim 4 wherein the proximal body section forms the soft second portion.
- A filter element as claimed in any of claims 1 to 4 wherein the filter body comprises a proximal body section and a distal body section interconnected by an intermediate body section, one or both of the proximal body section and the intermediate body section forming the soft second portion, the distal body section forming the stiff first portion.

7. A filter element as claimed in any of claims 4 to 6 wherein the proximal body section has a ribbed outer surface.

- 8. A filter element as claimed in any of claims 4 to 7 wherein a plurality of spaced-apart longitudinal ribs are provided on the proximal section.
- 9. A filter as claimed in any of claims 4 to 8 wherein the proximal body section includes corrugations.
- 20 10. A filter element substantially as hereinbefore described with reference to the accompanying drawings.

10

15

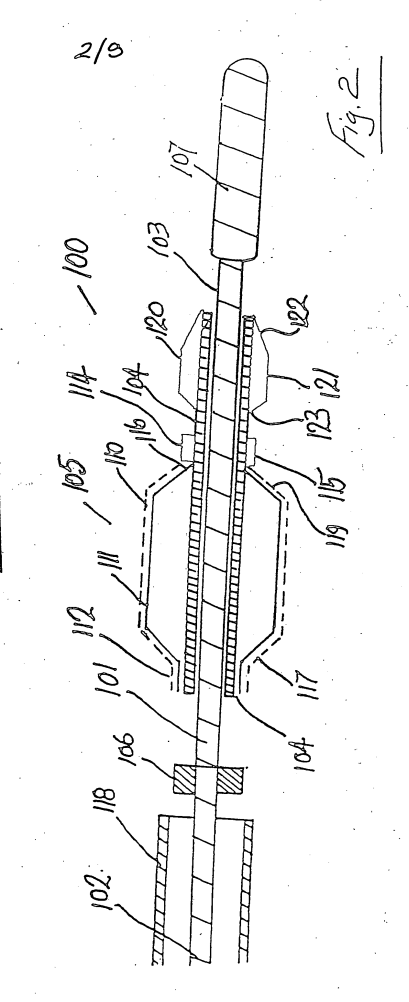
## **Abstract**

5

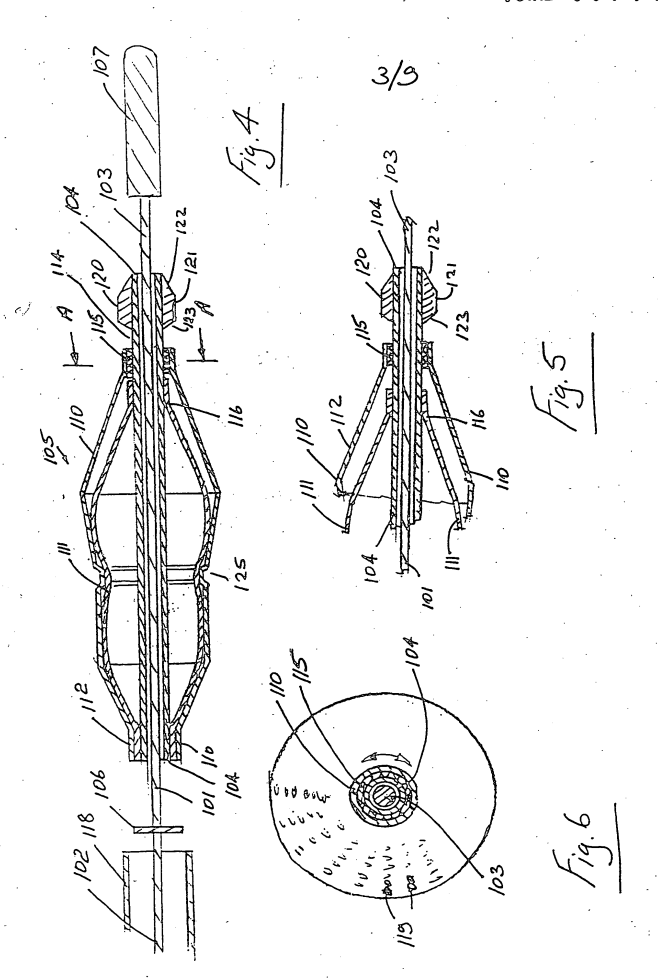
A collapsible filter element 2, 10, 20, 50, 40 for an embolic protection device comprises a filter body which has a relatively stiff portion 3 and a relatively soft portion 5 for engaging a vessel wall. The proximal and/or distal ends 3, 4 may be the stiff portion to provide enhanced web strength. To cater for a range of vessel sizes the filter body may have corrugations or ribs 46, 51.

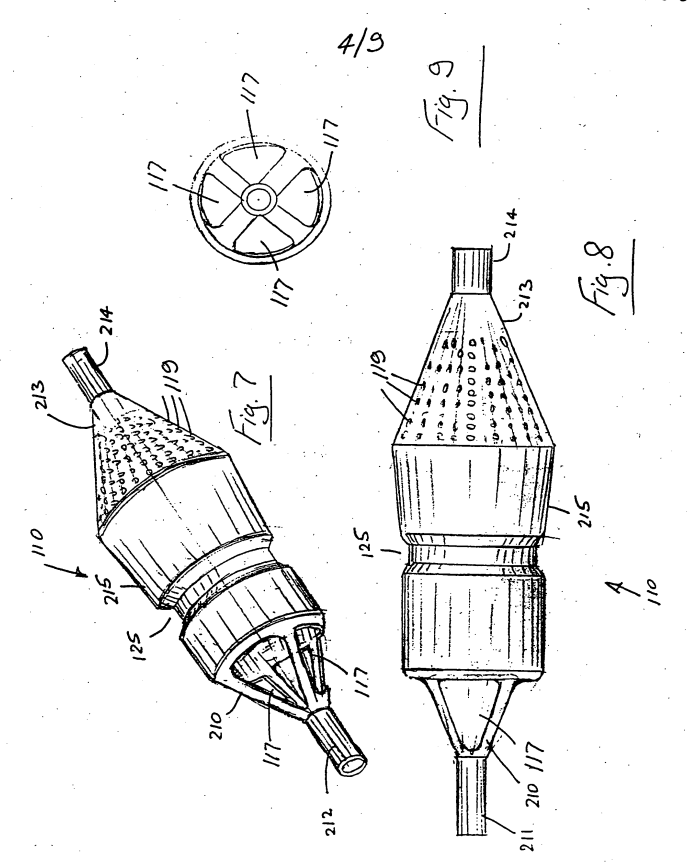
1/9

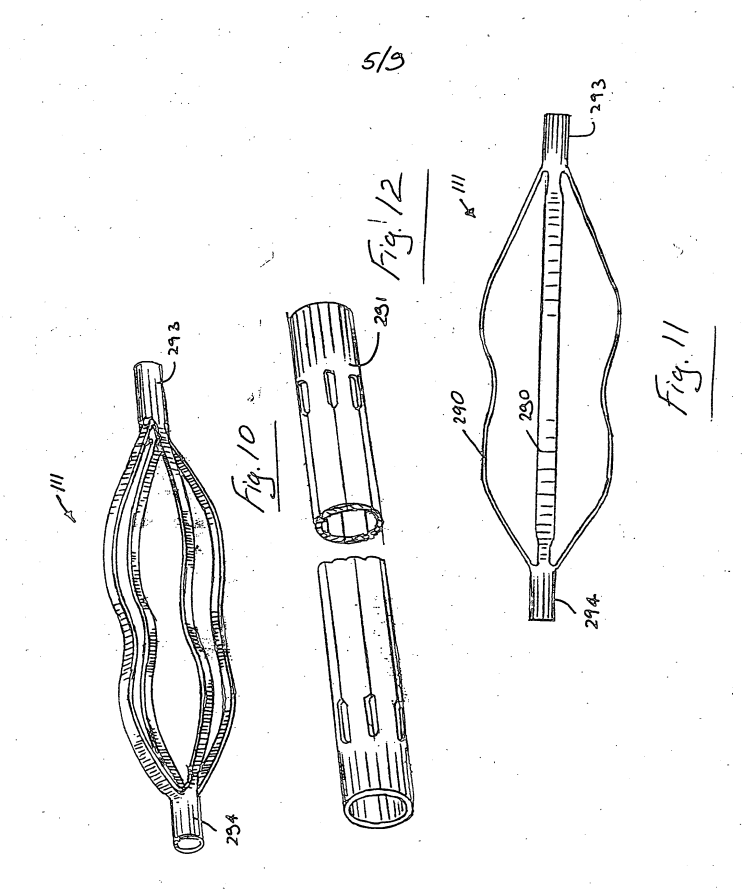
10/ 103



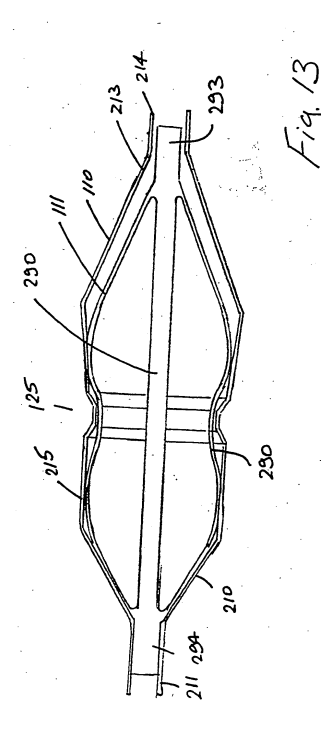
 $\subset$ 

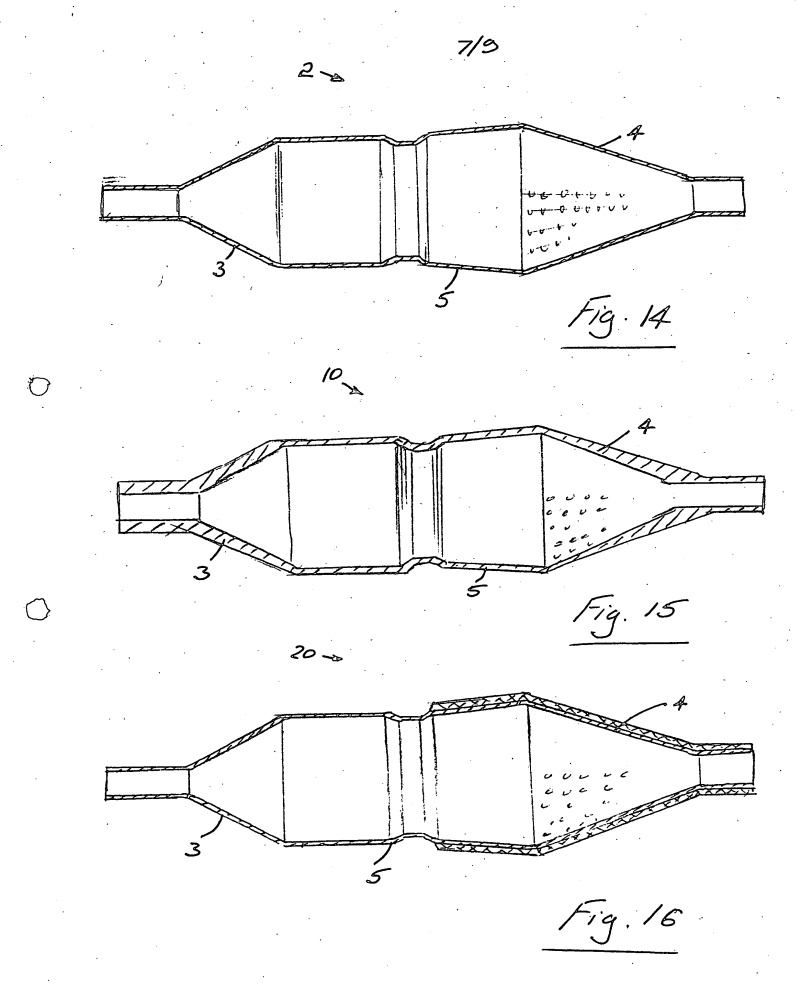






.





8/3

